Initial Approval: July 11, 2018

Revised Dated: January 8, 2020; April 10, 2019

CRITERIA FOR PRIOR AUTHORIZATION

Hepatitis C Agents

Provider Group Pharmacy

BILLING CODE TYPE For drug coverage and provider type information, see the KMAP Reference Codes webpage.

MANUAL GUIDELINES

All dosage forms of the medications listed in Table 1 below will require prior authorization.-Prior authorization will be required for all current and future dose forms available. All medication-specific criteria, including drug-specific indication, age, and dose for each agent is defined in table 1 below.

Elbasvir/grazoprevir (Zepatier®)

Glecaprevir/pibrentasvir (Mavyret®)

Ledipasvir/sofosbuvir (Harvoni®)

Ombitasvir/Paritaprevir/Ritonavir/Dasabuvir (Viekira Pak™)

Sofosbuvir (Sovaldi®)

Sofosbuvir/velpatasvir (Epclusa®)

Sofosbuvir/velpatasvir/voxilaprevir (Vosevi®)

CRITERIA FOR NON-REFRACTORY TREATMENT, INITIAL APPROVAL (MUST MEET ALL OF THE FOLLOWING):

*Patients new to the plan will be allowed to continue previous hepatitis C regimen (max of up to the duration listed below)

- Must be approved for the indication, age, genotype, and not exceed medication-specific quantity limit and duration of therapy listed in Table 1 and 2.¹⁻⁸
- Patient must have a diagnosis of chronic hepatitis C virus (HCV)
- Patient must have a confirmed genotype of 1a, 1b, 2, 3, 4, 5 or 6.
- For all agents listed, the preferred PDL drug, if applicable, which treats the PA indication, is required unless the patient meets the non-preferred PDL PA criteria.
- Patient has a pre-treatment <u>detectable</u> HCV RNA level drawn and results are submitted with PA request.
- Treatment regimen and duration of treatment must be prescribed in accordance with FDA-approved product labeling (defined in table 2)
- Requested medication must be prescribed for an FDA-approved age (defined in table 1)
- Dose must not exceed the medication-specific quantity limits (defined in table 1).
- Patient must not have a history of illicit intravenous (IV) substance use within the past 3 months.
- Prescriber must attest that the patient will be tested for evidence of current or prior hepatitis B virus (HBV) infection by measuring hepatitis B surface antigen (HBsAg) and hepatitis B core antibody (anti-HBc) before initiating HCV treatment.¹⁻⁸
- If the requested medication will be used in combination with ribavirin, female patients must have a negative pregnancy test within 30 days prior to initiation of therapy and monthly thereafter until treatment completion.
- Patient must not have been on previous or concurrent direct acting hepatitis C agents.
- Prescriber must attest that the patient's drug profile will be reviewed and monitored for potential clinically significant drug interactions (defined in table 1) with the requested medication prior to therapy initiation and throughout treatment duration.
- Prescriber must attest that all additional medication-specific safety criteria, as defined in table 1, is met.
- Prescriber must attest that the patient has been fully educated on their treatment and the importance of medication adherence and is motivated to be adherent to the full course of treatment.

• For all genotypes: the PDL preferred drug, which covers that specific genotype, is required unless the patient has a documented clinical rationale for using the non-preferred agent supported by the label.

LENGTH OF INITIAL APPROVAL: 4 weeks Up to the total number of approved weeks based upon FDA labeling in Table 2.

CRITERIA FOR RENEWAL FOR NON-REFRACTORY TREATMENT: (must meet all of the following)

Prescriber must document adherence by patient of greater than or equal to 90%.

LENGTH OF RENEWAL APPROVALS: 4 weeks, up to the total number of approved weeks based upon FDA labeling.

CRITERIA FOR REFRACTORYTREATMENT-EXPERIENCED (WITH PREVIOUS DAA) PATIENTS, INITIAL APPROVAL: (must meet all of the following)

- Patient must meet all criteria for non-refractorytreatment -, initial approval above.
- MCO claims data must indicate greater than or equal to 90% adherence to the previous direct-acting antiviral regimen (the MCO reviewer should verify this by the MCO claims data)
- Must be prescribed by or in consultation with a hepatologist, gastroenterologist, or infectious disease specialist
- The requested agent is FDA-approved as therapy for treatment-experienced patients. 1-8
- Patient has not been previously treated with and failed the requested regimen (regimen should include another DAA in which the patient has not failed).¹
- Prescriber has submitted documentation showing provided details that the patient has a documented presence of detectable HCV RNA at/up to least 12 weeks after the last completing treatment was given.¹
 - An assessment of viral response, including documentation of Sustained Viral Response (SVR), using an FDA-approved quantitative or qualitative nucleic acid test (NAT) with a detection level of greater than (>) 25 IU/mL at/up to 12 weeks after the last treatment was given (https://www.hcvguidelines.org/evaluate/when-whom)
- Prescriber has provided details that re-infection has been ruled out.
 - Patients who previously achieved SVR that have HCV recurrence due to reinfection may be managed as an initial infection.¹

LENGTH OF INITIAL APPROVAL: 4 weeks Up to the total number of approved weeks based upon FDA labeling in Table 2.

CRITERIA FOR REFRACTORY TREATMENT: (must meet all of the following)

Prescriber must document adherence by patient of greater than or equal to 90% for both agents.

LENGTH OF RENEWAL APPROVALS: 4 weeks, up to a total of 12 weeks based on approved treatment regimen and duration

FOR DRUGS THAT HAVE A CURRENT PA REQUIREMENT, BUT NOT FOR THE NEWLY APPROVED INDICATIONS, FOR OTHER FDA-APPROVED INDICATIONS, AND FOR CHANGES TO AGE REQUIREMENTS NOT LISTED WITHIN THE PA CRITERIA:

• THE PA REQUEST WILL BE REVIEWED BASED UPON THE FOLLOWING PACKAGE INSERT INFORMATION: INDICATION, AGE, DOSE, AND ANY PRE-REQUISITE TREATMENT REQUIREMENTS FOR THAT INDICATION.

<u>LENGTH OF APPROVAL (INITIAL AND RENEWAL): 8 weeks</u> Up to the total number of approved weeks based upon FDA labeling in the package insert.

TABLE 1: MEDICATION-SPECIFIC CRITERIA

MEDICATION	MEDICATION-SPECIFIC CRITERIA		
Daklinza®	Indication/Use	Hepatitis C virus (HCV) NS5A inhibitor indicated for use with sofosbuvir, with or without ribavirin,	
(daclatasvir)		for the treatment of chronic HCV genotype 1 or 3 infection.	
	Age (years)	<u>≥18</u>	
	Quantity Limit	1 tablet/day	
	Safety Criteria	➤ Patient must not be concurrently prescribed a strong CYP3A inducer (e.g. phenytoin,	
		carbamazepine, rifampin, St. John's wort)	
		→ Patient must not be on concurrent moderate CYP3A inducers (e.g. bosentan, dexamethasone,	
		efavirenz, etravirine, modafinil, nafcillin, rifapentine)	
Epclusa®	Indication/Use	Fixed dose combination of sofosbuvir, a hepatitis C virus (HCV) nucleotide analog NS5B	
(sofosbuvir/velpatasvir)	,	polymerase inhibitor, and velpatasvir, an HCV NS5A inhibitor, and is indicated for the treatment of	
		adult patients with chronic HCV genotype 1, 2, 3, 4, 5, or 6 infection:	
		without cirrhosis or with compensated cirrhosis	
		with decompensated cirrhosis for use in combination with ribavirin	
	Age (years)	<u>≥18</u>	
	Quantity Limit	1 tablet/day	
	Safety Criteria	→ Patient must not have severe renal impairment (eGFR < 30 mL/min/1.73m²) or currently require	
	,	hemodialysis	
		→ Patient must not be on concurrent:	
		• Amiodarone	
		 Moderate to strong inducers of CYP286 (e.g., carbamazepine, fosphenytoin, nevirapine, 	
		phenobarbital, phenytoin, primidone, rifampin)	
		 Moderate to strong inducers of CYP2C8 (e.g., rifampin) 	
		Moderate to strong inducers of CYP3A4 (e.g., avasimibe, carbamazepine, dexamethasone,	
		ethosuximide, griseofulvin, phenytoin, primidone, progesterone, rifabutin, rifampin, nafcillin, nelfinavir, nevirapine, oxcarbazepine, phenobarbital, phenylbutazone, St John's wort, sulfadimidine,	
		sulfinpyrazone, troglitazone)	
		 Inducers of P gp (e.g., avasimibe, carbamazepine, phenytoin, rifampin, St John's wort, 	
		tipranavir/ritonavir)	
Harvoni®	Indication/Use	Fixed dose combination of ledipasvir, a hepatitis C virus (HCV) NS5A inhibitor, and sofosbuvir, an	
(ledipasvir/sofosbuvir)		HCV nucleotide analog NS5B polymerase inhibitor, and is indicated for the treatment of chronic	
		hepatitis C virus (HCV) in:	
		 Adults with genotype 1, 4, 5, or 6 infection without cirrhosis or with compensated cirrhosis 	
		 Adults with genotype 1 infection with decompensated cirrhosis, in combination with 	
		ribavirin	
		 Adults with genotype 1 or 4 infection who are liver transplant recipients without cirrhosis 	
		or with compensated cirrhosis, in combination with ribavirin	
		 Pediatric patients 12 years of age and older or weighing at least 35 kg with genotype 1, 4, 	
		5, or 6 without cirrhosis or with compensated cirrhosis	
	Age (years)	≥ 12 years of age or weighing at least 35 kg	
	Quantity Limit	1 tablet/day	
	Safety Criteria	→ Patient must not have severe renal impairment (eGFR < 30 mL/min/1.73m²) or currently	
		require hemodialysis	
		→ If patient was on a previous course of treatment with Incivek or Victrelis it must have included	
		an interferon based regimen	
		→ Coadministration with amiodarone is not recommended. If alternative, viable treatment	
		options are unavailable, cardiac monitoring is recommended	
Mavyret®	Indication/Use	1. Fixed-dose combination of glecaprevir, a hepatitis C virus (HCV) NS3/4A protease inhibitor, and	
(glecaprevir/pibrentasvir)		pibrentasvir, an HCV NS5A inhibitor, and is indicated for the treatment of patients with chronic	
		HCV genotype (GT) 1, 2, 3, 4, 5 or 6 infection without cirrhosis and with compensated cirrhosis	
		(Child-Pugh A).	
		2. HCV genotype 1 infection, who previously have been treated with a regimen containing an HCV	
		NS5A inhibitor or an NS3/4A protease inhibitor, but not both.	
	Age (years)	<u>>18</u>	
	Quantity Limit	1 daily dose pack/day	
	Safety Criteria	→ Patient must not have moderate or severe hepatic impairment (Child-Pugh class B or C)	
		→—Patient must not be concurrently prescribed atazanavir or rifampin	
		→ Patient must not be on a concurrent direct acting hepatitis C agent or ribavirin	

TABLE 1 (CONT.). MEDICATION-SPECIFIC CRITERIA

MEDICATION	MEDICATION-SPECIFIC CRITERIA		
Olysio® (simeprevir)	Indication/Use	Hepatitis C virus (HCV) NS3/4A protease inhibitor indicated for the treatment of adults with chronic hepatitis C virus (HCV) infection: - in combination with sofosbuvir in patients with HCV genotype 1 without cirrhosis or with compensated cirrhosis - in combination with peginterferon alfa (Peg IFN alfa) and ribavirin (RBV) in patients with HCV genotype 1 or 4 without cirrhosis or with compensated cirrhosis	
	Age (years)	<u>>18</u>	
	Quantity Limit	1 capsule/day	
	Safety Criteria	→ If patient has subtype 1a they must have a negative test for NS3-Q80k polymorphism	
	Sarety enteria	The patient must not have advanced and/or decompensated cirrhosis (moderate or severe hepatic impairment)	
Sovaldi® (sofosbuvir)	Indication/Use	Hepatitis C virus (HCV) nucleotide analog NS5B polymerase inhibitor indicated for the treatment of:	
		 Adult patients with genotype 1, 2, 3 or 4 chronic hepatitis C virus (HCV) infection without cirrhosis or with compensated cirrhosis as a component of a combination antiviral treatment regimen. Pediatric patients 12 years of age and older or weighing at least 35 kg with genotype 2 or chronic HCV infection without cirrhosis or with compensated cirrhosis in combination with 	
	A = = ()	ribavirin.	
	Age (years)	≥-18 (genotype 1, 2, 3, 4)	
	0 111 11 11	≥12 years of age or weighing at least 35 kg (genotype 2 or 3)	
	Quantity Limit	1 tablet/day	
	Safety Criteria	→ Coadministration with amiodarone is not recommended. If alternative, viable treatment options are unavailable, cardiac monitoring is recommended	
Technivie®	Indication/Use	Fixed-dose combination of ombitasvir, a hepatitis C virus NSSA inhibitor, paritaprevir, a hepatitis C	
(ombitasvir/paritaprevir/ ritonavir)	mulcation, ose	virus NS3/4A protease inhibitor, and ritonavir, a CYP3A inhibitor and is indicated in combination with ribavirin for the treatment of patients with genotype 4 chronic hepatitis C virus (HCV) infection without cirrhosis or with compensated cirrhosis.	
	Age (years)	→ 18 —	
	Quantity Limit	2 tablets/day	
	Safety Criteria	→ Patient must not have moderate or severe hepatic impairment or cirrhosis (Metavir score of F4 and Child-Pugh class B or C)	
		→ Patient must not be concurrently prescribed a moderate or strong CYP3A inducer	
Viekira Pak™, Viekira XR™ (ombitasvir/paritaprevir/ ritonavir and dasabuvir)	Indication/Use	Treatment of adult patients with chronic hepatitis C virus (HCV): - genotype 1b without cirrhosis or with compensated cirrhosis - genotype 1a without cirrhosis or with compensated cirrhosis for use in combination with ribavirin. (VIEKIRA PAK includes ombitasvir, a HCV NS5A inhibitor, paritaprevir, a HCV NS3/4A protease inhibitor,	
	A == (++=====)	ritonavir, a CYP3A inhibitor and dasabuvir, a HCV non-nucleoside NS5B palm polymerase inhibitor)	
	Age (years) Quantity Limit	≥ 18 1 daily dose pack/day	
	Safety Criteria	→ Patient must not have underlying moderate to severe hepatic impairment (Child-Pugh class B or C)	
Zepatier® (elbasvir/grazoprevir)	Indication/Use	Fixed dose combination product containing elbasvir, a hepatitis C virus (HCV) NS5A inhibitor, and grazoprevir, an HCV NS3/4A protease inhibitor, and is indicated for treatment of chronic HCV genotype 1 or 4 infection in adults. ZEPATIER is indicated for use with ribavirin in certain patient populations.	
	Age (years)	<u>≥18</u>	
	Quantity Limit	1 tablet/day	
	Safety Criteria	 Patient must not be concurrently prescribed a strong CYP3A inducer (e.g. phenytoin, carbamazepine, rifampin, St. John's Wort), efavirenz, or OATP1B1/3 inhibitor (e.g. cyclosporine eltrombopag, lapatinib, lopinavir, rifampin, ritonavir) If the patient has genotype 1a, patient must be tested for the presence of virus with NS5A resistance-associated polymorphisms prior to initiation of therapy 	
		→ Patient must not have moderate or severe hepatic impairment (Child-Pugh class B or C)	

TABLE 1 (CONT.). MEDICATION-SPECIFIC CRITERIA

MEDICATION	MEDICATION-SPECIFIC CRITERIA	
Vosevi™ (sofosbuvir/velpatasvir/ voxilaprevir)	Indication/Use	Fixed-dose combination of sofosbuvir, a hepatitis C virus (HCV) nucleotide analog NS5B polymerase inhibitor, velpatasvir, an HCV NS5A inhibitor, and voxilaprevir, an HCV NS3/4A protease inhibitor, and is indicated for the treatment of adult patients with chronic HCV infection without cirrhosis or with compensated cirrhosis (Child-Pugh A) who have: - Genotype 1, 2, 3, 4, 5, or 6 infection and have previously been treated with an HCV regimen containing an NS5A inhibitor Genotype 1a or 3 infection and have previously been treated with an HCV regimen containing sofosbuvir without an NS5A inhibitor.
	Age (years) Quantity Limit Safety Criteria	≥ 18 1 tablet/day → Patient must not have severe renal impairment (eGFR < 30 mL/min/1.73m²) or currently require hemodialysis → Patient must not be on concurrent rifampin
		→ Patient should not be on concurrent: P-gp inducers, moderate to potent CYP2B6, 2C8, or 3A4 inducers, amiodarone (if alternative, viable treatment options are unavailable, cardiac monitoring is recommended)

TABLE 2. TREATMENT REGIMEN AND DURATION

MEDICATION	GENOTYPE	PATIENT POPULATION	TREATMENT AND DURATION
Daklinza®		Without cirrhosis	Daklinza + Sofosbuvir for 12 weeks
(daclatasvir)		Compensated (Child Pugh A) cirrhosis	Daklinza + Sofosbuvir for 12 weeks
	1	Decompensated (Child-Pugh B or C) cirrhosis	Daklinza + Sofosbuvir + Ribavirin for 12 weeks
		Post-transplant	Daklinza + Sofosbuvir + Ribavirin for 12 weeks
		Without cirrhosis	Daklinza + Sofosbuvir for 12 weeks
		Compensated (Child Pugh A) cirrhosis	Daklinza + Sofosbuvir + Ribavirin for 12 weeks
	3	Decompensated (Child-Pugh B or C) cirrhosis	Daklinza + Sofosbuvir + Ribavirin for 12 weeks
		Post-transplant	Daklinza + Sofosbuvir + Ribavirin for 12 weeks
Epclusa® (sofosbuvir/velpatasvir)		Treatment-naïve and treatment-experienced*, without cirrhosis and with compensated cirrhosis (Child-Pugh A)	Epclusa for 12 weeks
	1, 2, 3, 4, 3, 6	Treatment naïve and treatment experienced ^a ; with decompensated cirrhosis (Child-Pugh B or C)	Epclusa + Ribavirin for 12 weeks
Harvoni® (ledipasvir/sofosbuvir)		Treatment-naïve without cirrhosis or with compensated cirrhosis (Child Pugh A)	Harvoni for 12 weeks
		Treatment-experienced ^b -without cirrhosis	Harvoni for 12 weeks
	1	Treatment-experienced ^b -with compensated cirrhosis (Child-Pugh A)	Harvoni for 24 weeks
		Treatment-naïve and treatment-experienced ^b with decompensated cirrhosis (Child-Pugh B or C)	Harvoni + Ribavirin for 12 weeks
	1 or 4	Treatment-naïve and treatment-experienced ^b liver transplant recipients without cirrhosis, or with compensated cirrhosis (Child-Pugh A)	Harvoni + Ribavirin for 12 weeks
	4, 5 or 6	Treatment-naïve and treatment-experienced ^b , without cirrhosis and with compensated cirrhosis (Child-Pugh A)	Harvoni for 12 weeks

TABLE 2 (CONT.). TREATMENT REGIMEN AND DURATION

MEDICATION	GENOTYPE	PATIENT POPULATION	TREATMENT AND DURATION
Mavyret® (glecaprevir/pibrentasvir)	1, 2, 3, 4, 5, 6	Treatment-naïve, without cirrhosis	Mavyret for 8 weeks
		Treatment-naïve, with compensated cirrhosis (Child-Pugh A)	Mavyret for 12 weeks
	4	Treatment experienced, previously treated with regimen containing an NS5A inhibitor ^c without prior treatment with an NS3/4A protease inhibitor, without cirrhosis, or with compensated cirrhosis (Child-Pugh A)	Mavyret for 16 weeks
	1	Treatment experienced, previously treated with regimen containing an NS3/4A PI ^d without prior treatment with an NS5A inhibitor, without cirrhosis, or with compensated cirrhosis (Child-Pugh A)	Mavyret for 12 weeks
		Treatment-experienced, previously treated with regimen containing PRSe, without cirrhosis	Mavyret for 8 weeks
	1, 2, 4, 5, 6	Treatment-experienced, previously treated with regimen containing PRSe, with compensated cirrhosis (Child-Pugh A)	Mavyret for 12 weeks
	3	Treatment experienced, previously treated with regimen containing PRS ^e , without cirrhosis, or with compensated cirrhosis (Child-Pugh A)	Mavyret for 16 weeks
Olysio® (simeprevir)	1	Treatment-naïve and treatment-experienced, without cirrhosis	Olysio + Sofosbuvir for 12 weeks
(1	Treatment-naïve and treatment-experienced, with compensated cirrhosis (Child Pugh A)	Olysio + Sofoxbuvir for 24 weeks
	1, 4	Treatment-naïve and treatment-experienced, without cirrhosis or with compensated cirrhosis (Child-Pugh A), with or without HIV-1 co- infection	Olysio + Peg-IFN-alfa + Ribavirin for 12 weeks (followed by 12 or 36 additional weeks of Peg-IFN alfa + Ribavirin depending on prior response status and presence of HIV 1 infection)
Sovaldi® (sofosbuvir)	1, 4	Treatment naïve without cirrhosis or with compensated cirrhosis (Child Pugh A)	Sovaldi + Peg-IFN-alfa + Ribavirin for 12 weeks
(3010354411)	2	Treatment-naïve and treatment-experienced ^b without cirrhosis or with compensated cirrhosis (Child-Pugh A)	Sovaldi + Ribavirin for 12 weeks
	3	Treatment naïve and treatment experiencedb without cirrhosis or with compensated cirrhosis (Child-Pugh A)	Sovaldi + Ribavirin for 24 weeks
Technivie® (ombitasvir/paritaprevir/ ritonavir)	4	Without cirrhosis or with compensated cirrhosis (Child-Pugh A)	Technivie + Ribavirin for 12 weeks (Technivie without ribavirin for 12 weeks may be considered for treatment naïve patients without cirrhosis who cannot take or tolerate ribavirin)
Viekira Pak™, Viekira XR™		Treatment-naïve or interferon-experienced, without cirrhosis	Viekira Pak + Ribavirin for 12 weeks
(ombitasvir/paritaprevir/ ritonavir and dasabuvir)	1a	Treatment-naïve or interferon-experienced, with compensated cirrhosis (Child-Pugh A)	Viekira Pak + Ribavirin for 24 weeks (Viekira Pak + ribavirin for 12 week may be considered for some patients based on prior treatment history)
	1b	Treatment-naïve or interferon-experienced, without cirrhosis or with compensated cirrhosis	Viekira Pak for 12 weeks

TABLE 2 (CONT.). TREATMENT REGIMEN AND DURATION

MEDICATION	GENOTYPE	PATIENT POPULATION	TREATMENT AND DURATION
Zepatier®		Treatment-naïve or Peg-IFN/Ribavirin-	
(elbasvir/grazoprevir)	1a	experienced without baseline NS5A	Zepatier for 12 weeks
, , , , , ,	10	polymorphisms, without cirrhosis, or with	Zepatier for 12 weeks
		compensated cirrhosis (Child-Pugh A)	
		Treatment naïve or Peg IFN/Ribavirin	
		experienced with baseline NS5A	Zepatier + Ribavirin for 16 weeks
		polymorphisms, without cirrhosis, or with	zepatier + Kibavirin for 16 weeks
		compensated cirrhosis (Child Pugh A)	
		Treatment-naïve or Peg-IFN/Ribavirin-	
	1b	experienced without cirrhosis, or with	Zepatier for 12 weeks
		compensated cirrhosis (Child-Pugh A)	
		Peg IFN/Ribavirin/NS3/4A protease inhibitor	
	1a or 1b	experienced, without cirrhosis, or with	Zepatier + Ribavirin for 12 weeks
		compensated cirrhosis (Child-Pugh A)	
		Treatment naïve, without cirrhosis, or with	Zepatier for 12 weeks
		compensated cirrhosis (Child-Pugh A)	zepatier for 12 weeks
	4	Peg-IFN/Ribavirin/NS3/4A protease inhibitor-	
		experienced, without cirrhosis, or with	Zepatier + Ribavirin for 16 weeks
		compensated cirrhosis (Child-Pugh A)	
Vosevi™		Treatment experienced, previously treated with	
(sofosbuvir/velpatasvir/	4 2 2 4 5 6	regimen containing an NS5∆ inhibitor ^g , without	V :6 40 1
voxilaprevir)	1, 2, 3, 4, 5, 6	cirrhosis or with compensated cirrhosis (Child-	Vosevi for 12 weeks
voxilaprevii j		Pugh A)	
		Treatment-experienced, previously treated with	
		regimen containing sofosbuvir without an NS5A	Wasself for 42 weeks
	1a or 3	inhibitorh, without cirrhosis or with	Vosevi for 12 weeks
		compensated cirrhosis (Child-Pugh A)	

a—In clinical trials, regimens contained peginterferon alfa/ribavirin with or without an HCV NS3/4A protease inhibitor (boceprevir, simeprevir, or telaprevir)

Table 1. FDA-approved age and indications for Hepatitis C Agents.²⁻⁸

<u>Agents</u>	Indication(s)	Age/Weight		
<u>Antihepacivira</u>	NS3/4A Protease Inhibitor and NS5A Inhibitor Combination			
Elbasvir/Grazoprevir (Zepatier®)	Chronic hepatitis C genotype 1 or 4 infection without	≥ 18 years		
	cirrhosis or with compensated cirrhosis (Child-Pugh A)			
Glecaprevir/pibrentasvir	Chronic hepatitis C genotype 1, 2, 3, 4, 5, or 6 infection	≥ 12 years or		
(Mavyret®)	without cirrhosis or with compensated cirrhosis (Child-Pugh	weighing ≥ 45 kg		
	<u>A)</u>			
Antihepaciviral NS3/4A I	Antihepaciviral NS3/4A Protease Inhibitor and NS5A Inhibitor and NS5B Inhibitor Combination			
Ombitasvir/Paritaprevir/	Chronic hepatitis C genotype 1a or 1b infection without	≥ 18 years		
Ritonavir/Dasabuvir (Viekira Pak™)	onavir/Dasabuvir (Viekira Pak™) cirrhosis or with compensated cirrhosis (Child-Pugh A)			
Sofosbuvir/velpatasvir/	Chronic hepatitis C genotype 1, 2, 3, 4, 5, or 6 infection	≥ 18 years		
voxilaprevir (Vosevi®)	voxilaprevir (Vosevi®) without cirrhosis or with compensated cirrhosis (Child-Pugh			
<u>A)</u>				
Antihepaciviral NS5A Inhibitor and NS5B Inhibitor Combination				
<u>Ledipasvir/sofosbuvir (Harvoni®)</u>	Chronic hepatitis C genotype 1, 4, 5, or 6 infection	≥ 3 years		

b—Treatment experienced patients have failed an interferon based regimen with or without ribavirin

e-In clinical trials, subjects were treated with prior regimens containing ledipasvir and sofosbuvir or daclatasvir with pegylated interferon and ribavirin

^d—In clinical trials, subjects were treated with prior regimens containing simeprevir and sofosbuvir, or simeprevir, boceprevir, or telaprevir with pegylated interferon and ribavirin

e—PRS = Prior treatment experience with regimens containing interferon, pegylated interferon, ribavirin, and/or sofosbuvir, but no prior treatment experience with an HCV NS3/4A PL or NS5A inhibitor

f—Treatment experienced patients include prior relapsers, prior partial responders and prior null responders who failed prior IFN based therapy

⁶_—In clinical trials, prior NSSA inhibitor experience included daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir

h—In clinical trials, prior treatment experience included sofosbuvir with or without any of the following: peginterferon alfa/ribavirin, ribavirin, HCV NS3/4A protease inhibitor (boceprevir, simeprevir, or telaprevir)

Sofosbuvir/Velpatasvir (Epclusa®)	Chronic hepatitis C genotype 1, 2, 3, 4, 5, or 6 infection	≥ 18 years
	Antihepaciviral NS5B Inhibitor	
Sofosbuvir (Sovaldi®)	Chronic hepatitis C genotype 1, 2, 3, or 4 infection in adults without cirrhosis or with compensated cirrhosis (Child-Pugh A) as a component of a combination antiviral treatment regimen.	≥ 18 years
	Chronic hepatitis C genotype 2 or 3 infection in pediatrics without cirrhosis or with compensated cirrhosis (Child-Pugh A) in combination with ribavirin.	≥ 3 years

Table 2. Treatment Regimen and Duration by Genotype.²⁻⁸

Agents	Patient Population	<u>Treatment Duration</u>
Antil	nepaciviral NS3/4A Protease Inhibitor and NS5A Inhibitor Co	ombination
Elbasvir/Grazoprevir	Genotype 1a and treatment-naïve or	One tablet daily (elbasvir 50
(Zepatier®)	peginterferon/ribavirin-experienced without cirrhosis or	mg-grazoprevir 100 mg per
	with compensated cirrhosis (Child-Pugh class A) without	day) for 12 weeks.
	baseline NS5A polymorphisms (at amino acid positions	
	28, 30, 31, or 93).	
	Genotype 1b and treatment-naïve or	
	peginterferon/ribavirin-experienced without cirrhosis or	
	with compensated cirrhosis (Child-Pugh class A).	
	Genotype 4 and treatment-naïve without cirrhosis or with	
	compensated cirrhosis (Child-Pugh class A).	
	Genotype 1a or 1b and treatment-experienced with	One tablet daily (elbasvir 50
	peginterferon/ribavirin/HCV NS3/4A protease inhibitor	mg-grazoprevir 100 mg per
	without cirrhosis or with compensated cirrhosis (Child-	day) for 12 weeks in
	Pugh class A).	combination with ribavirin.
	Genotype 1a and treatment-naïve or	One tablet daily (elbasvir 50
	peginterferon/ribavirin-experienced without cirrhosis or	mg-grazoprevir 100 mg per
	with compensated cirrhosis (Child-Pugh class A) with	day) for 16 weeks in
	baseline NS5A polymorphisms (at amino acid positions	combination with ribavirin.
	28, 30, 31, or 93).	
	Genotype 4 and treatment-experienced with	
	peginterferon/ribavirin without cirrhosis or with	
	compensated cirrhosis (Child-Pugh class A).	
Glecaprevir/pibrentasvir	Genotype 1, 2, 3, 4, 5, 6, and treatment-naïve without	Three tablets daily
(Mavyret®)	<u>cirrhosis or with compensated cirrhosis (Child-Pugh class</u>	(glecaprevir 300 mg-
	<u>A).</u>	pibrentasvir 120 mg per day)
		for 8 weeks.
	Genotype 1, 2, 4, 5, 6, and treatment-experienced with	
	peginterferon/ribavirin and/or sofosobuvir (without prior	

DRAFT FA CITTETIA	treatment with an NS5A inhibitor or NS3/4A protease	
	inhibitor) without cirrhosis.	
	Genotype 1 and treatment-experienced with an NS3/4A	Three tablets daily
	protease inhibitor (without prior treatment with an NSSA	(glecaprevir 300 mg-
	inhibitor) without cirrhosis or with compensated cirrhosis	pibrentasvir 120 mg per day)
	(Child-Pugh class A).	for 12 weeks.
	Genotype 1, 2, 4, 5, 6, and treatment-experienced with	
	peginterferon/ribavirin and/or sofosobuvir (without prior	
	treatment with an NS5A inhibitor or NS3/4A protease	
	inhibitor) with compensated cirrhosis (Child-Pugh class	
	<u>A).</u>	
	Genotype 1, 2, 4, 5, 6, and liver or kidney transplant	
	recipients without cirrhosis or with compensated	
	cirrhosis (Child-Pugh class A).	
	Genotype 1 and treatment-experienced with an NS5A	Three tablets daily
	inhibitor (without prior treatment with an NS3/4A	(glecaprevir 300 mg-
	protease inhibitor) without cirrhosis or with compensated	pibrentasvir 120 mg per day)
	cirrhosis (Child-Pugh class A).	for 16 weeks.
	CHITTOSIS (CHITA T Agri Class 74).	101 10 WCCR3.
	Genotype 3 and treatment-experienced with	
	peginterferon/ribavirin and/or sofosobuvir ((without	
	prior treatment with an NS5A inhibitor or NS3/4A	
	protease inhibitor) without cirrhosis or with compensated	
	cirrhosis (Child-Pugh class A).	
	Genotype 1 and liver or kidney transplant recipient's	
	treatment-experienced with an NS5A inhibitor (without	
	prior treatment with an NS3/4A protease inhibitor)	
	without cirrhosis or with compensated cirrhosis (Child-	
	Pugh class A).	
	<u> </u>	
	Genotype 3 and liver or kidney transplant recipient's	
	treatment-experienced with peginterferon/ribavirin	
	and/or sofosobuvir (without prior treatment with an	
	NS5A inhibitor or NS3/4A protease inhibitor) with	
	compensated cirrhosis (Child-Pugh class A).	
Antihepacivira	I NS3/4A Protease Inhibitor and NS5A Inhibitor and NS5B Ir	hhibitor Combination
Ombitasvir/Paritaprevir/	Genotype 1a without cirrhosis	Four tablets daily (ombitasvir
Ritonavir/Dasabuvir		25 mg-paritaprevir 150 mg-
(Viekira Pak™)		ritonavir 100 mg-dasabuvir
		500 mg per day) with
		concomitant ribavirin for 12
		weeks.
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	Genotype 1a with compensated cirrhosis	Four tablets daily (ombitasvir
		25 mg-paritaprevir 150 mg-
		ritonavir 100 mg-dasabuvir
		500 mg per day) with
		concomitant ribavirin for 24
		weeks.
		* Medication administered
		with ribavirin for 12 weeks
		may be
		considered for patients with
		prior pegIFN and who partially
		responded.
	Genotype 1b without cirrhosis or with compensated	Four tablets daily (ombitasvir
	<u>cirrhosis</u>	25 mg-paritaprevir 150 mg-
		ritonavir 100 mg-dasabuvir
		500 mg per day) for 12 weeks.
Sofosbuvir/velpatasvir/	Genotype 1, 2, 3, 4, 5, 6, and treatment-experienced with	One tablet daily (sofosbuvir
voxilaprevir (Vosevi®)	an NS5A inhibitor without cirrhosis or with compensated	400 mg-valpatasvir 100 mg-
	cirrhosis (Child-Pugh class A).	voxilaprevir 100 mg per day)
		for 12 weeks.
	Genotype 1a or 3, and treatment-experienced with	
	sofosbuvir (without prior treatment with an NSSA	
	inhibitor) without cirrhosis or with compensated cirrhosis	
	(Child-Pugh class A).	
	Antihepaciviral NS5A Inhibitor and NS5B Inhibitor Combin	ation
Ledipasvir/sofosbuvir	Genotype 1 and treatment-naïve without cirrhosis or with	Pediatrics weighing ≥ 35 kg
(Harvoni®)	compensated cirrhosis (Child-Pugh class A).	and adults: one tablet or
(Harvoin)	compensated cirriosis (cinia i agriciass A).	packet daily (ledipasvir 90 mg-
	Genotype 1 and treatment-experienced without cirrhosis.	sofosbuvir 400 mg per day) for
		12 weeks.
	Genotype 4, 5, 6, and treatment-naïve or treatment-	
	experienced without cirrhosis or with compensated	Pediatrics weighing 17 to < 35
	cirrhosis (Child-Pugh class A).	kg: one tablet or packet daily
		(ledipasvir 45 mg-sofosbuvir
		200 mg per day) for 12 weeks.
		Pediatrics weighing < 17 kg:
		one tablet or packet daily
		(ledipasvir 33.75 mg-
		sofosbuvir 150 mg per day) for
		12 weeks.
	Genotype 1 and treatment-naïve or treatment-	Pediatrics weighing ≥ 35 kg
	experienced with decompensated cirrhosis (Child-Pugh	and adults: one tablet or
	class B or C).	packet daily (ledipasvir 90 mg-
		sofosbuvir 400 mg per day)

Genotype 1 or 4, and treatment-naïve or treatment-	with concomitant ribavirin for
	12 weeks.
	12 WEEKS.
or with compensated cirriosis (child-Pugh class A).	Dedictrics weighing 17 to 425
	Pediatrics weighing 17 to < 35
	kg: one tablet or packet daily
	(ledipasvir 45 mg-sofosbuvir
	200 mg per day) with
	concomitant ribavirin for 12
	weeks.
	Pediatrics weighing < 17 kg:
	one tablet or packet daily
	(ledipasvir 33.75 mg-
	sofosbuvir 150 mg per day)
	with concomitant ribavirin for
	12 weeks.
	Pediatrics weighing ≥ 35 kg
compensated cirrhosis (Child-Pugh class A).	and adults: one tablet or
	packet daily (ledipasvir 90 mg-
	sofosbuvir 400 mg per day) for
	24 weeks.
	Pediatrics weighing 17 to < 35
	kg: one tablet or packet daily
	(ledipasvir 45 mg-sofosbuvir
	200 mg per day) for 24 weeks.
	200 mg per day) for 24 weeks.
	Pediatrics weighing < 17 kg:
	one tablet or packet daily
	(ledipasvir 33.75 mg-
	sofosbuvir 150 mg per day) for
	24 weeks.
Genotype 1, 2, 3, 4, 5, 6, and treatment-naïve or	One tablet daily (sofosbuvir
peginterferon/ribavirin-experienced with or without an	400mg-velpatasvir 100mg per
HCV NS3/4A protease inhibitor (boceprevir, simeprevir,	day) for 12 weeks.
or telaprevir) without cirrhosis or with compensated	
cirrhosis (Child-Pugh class A).	
Genotype 1, 2, 3, 4, 5, 6, and treatment-naïve and	One tablet daily (sofosbuvir
treatment-experienced with or without an HCV NS3/4A	400mg-velpatasvir 100mg per
protease inhibitor	day) with concomitant
with decompensated cirrhosis (Child-Pugh B and C).	ribavirin for 12 weeks.
Antihepaciviral NS5B Inhibitor	
Adults and pediatrics with genotype 2 and treatment-	Pediatrics weighing ≥ 35 kg
1	1
naïve or treatment-experienced without cirrhosis or with	and adults: One tablet or
	Genotype 1 and treatment-experienced with compensated cirrhosis (Child-Pugh class A). Genotype 1 and treatment-experienced with compensated cirrhosis (Child-Pugh class A). Genotype 1, 2, 3, 4, 5, 6, and treatment-naïve or peginterferon/ribavirin-experienced with or without an HCV NS3/4A protease inhibitor (boceprevir, simeprevir, or telaprevir) without cirrhosis or with compensated cirrhosis (Child-Pugh class A). Genotype 1, 2, 3, 4, 5, 6, and treatment-naïve and treatment-experienced with or without an HCV NS3/4A protease inhibitor without an HCV NS3/4A protease inhibitor without an HCV NS3/4A protease inhibitor with decompensated cirrhosis (Child-Pugh B and C). Antihepaciviral NS5B Inhibitor

One tablet or packet daily
(sofosbuvir 150 mg per day)
with concomitant
peginterferon and ribavirin for
12 weeks.

Adults and pediatrics with genotype 3 and treatmentnaïve or treatment-experienced without cirrhosis or with
compensated cirrhosis (Child-Pugh class A).

Pediatrics weighing ≥ 35 kg
and adults: One tablet or
packet daily (sofosbuvir 400
mg per day) with concomitant
ribavirin for 24 weeks.

Pediatrics weighing 17 to < 35
kg: One tablet or packet daily
(sofosbuvir 200 mg per day)
with concomitant ribavirin for
24 weeks.

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	Pediatrics weighing < 17 kg: One tablet or packet daily (sofosbuvir 150 mg per day) with concomitant ribavirin f 24 weeks.
· · · · · · · · · · · · · · · · · · ·	rir) was discontinued by Abbvie in January 2019. rir/dasabuvir) was discontinued by Abbvie in January 2019. rissen in May 2018. Merck in December 2015.
 Epclusa (sofosbuvir/velpatasvir) [prescribin 2019. Harvoni (ledipasvir/sofosbuvir) [prescribing 2019. Mavyret (glecaprevir/pibrentasvir) [prescribing information of the second of the seco	g, managing, and treating hepatitis C. November 30, 2019. Available at: https://www.hcvguidelines.org/ ng information]. Foster City, CA: Gilead Sciences, Inc; November g information]. Foster City, CA: Gilead Sciences, Inc; November ibing information]. North Chicago, IL: AbbVie Inc; September 2019. on]. Foster City, CA: Gilead Sciences, Inc; September 2019. avir/dasabuvir) [prescribing information]. North Chicago, IL: AbbVie r) [prescribing information]. Foster City, CA: Gilead Sciences Inc; ibing information]. Whitehouse Station, NJ: Merck Sharp & Dohme
DRUG UTILIZATION REVIEW COMMITTEE CHAIR	PHARMACY PROGRAM MANAGER DIVISION OF HEALTH CARE FINANCE

DATE

DATE

KANSAS DEPARTMENT OF HEALTH AND ENVIRONMENT